

National Library of Medicine

PubMed

PubMed	Nucleotide	Protein	Genome	Structure	PopSet	Taxonomy	OMIM
Search Publ	Med ▼ for			The second secon	Many on and contrates a contrate Mile	Go Cl	ear
Cycles Cas American	and the second of the second o	Limits	Preview/Ind	lex Histo	ory	Clipboard	
25.7	Dis	splay Abstract	a sucure transment april account of the contract of the contra	Save Text	Order	Add to Clipbo	ard
	3 1	Mount Con	m Ganamics	1998:3(3) 173	7 <u>-</u> 92	the selection and	bués pouns

 $\equiv 1$. Microb (omp Genomics 1998:5(5) 177-9 $_{-}$

Constructing multigenome views of whole microbial genomes.

Gaasterland T, Ragan MA

Mathematics and Computer Science Division, Argonne National Labortory, Illinois, USA.

We have designed and implemented a system to carry out cross-genome comparisons of open reading frames (ORFs) from multiple genomes. This implementation includes a genome profiling system that allows us to explore pairwise comparisons at different levels of match similarity and ask biologically motivated queries involving number and identity of ORFs, their function, functional category, distribution in genomes or in biological domains, and statistics on their matches and match families. This analysis required precise definition of new classification terms and concepts. We define the terms genomic signature, summary signature, biologic domain signature, domain class, match level, match family, and extended match family, then use these terms to define concepts, including genomically universal proteins and proteins characteristics of sets of genomes. We initiate an analysis based on automated FASTA (Pearson, 1996) comparison of 22.419 conceptually translated protein sequences from nine microbial genomes.

PMID: 9775388

Display	Abstract	▼ Save Text	Order	Add to Clipboard

Write to the Help Desk NCBI | NLM | NIH Department of Health & Human Services Freedom of Information Act | Disclaimer